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# UniProtKB/TrEMBL entry Q9XEI0



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Entry history

## [Entry info] [Name and origin] [References] [Comments] [Cross-references] [Keywords] [Features] [Sequence] [Tools]

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## **Entry information**

Entry name

**Q9XEIO STERE** 

Primary accession number

Q9XEI0

None

Secondary accession numbers Integrated into TrEMBL on

November 1, 1999

Sequence was last modified on

November 1, 1999 (Sequence version 1)

Annotations were last modified on

April 18, 2006 (Entry version 23)

## Name and origin of the protein

Protein name

Kaurene synthase

Synonyms

None

Gene name

Name: KS22-1

From Taxonomy Stevia rebaudiana (Stevia) [TaxID: 55670]

Eukaryota; Viridiplantae; Streptophyta; Embryophyta;

Tracheophyta; Spermatophyta; Magnoliophyta;

eudicotyledons: core eudicotyledons: asterids: campanu

Asterales; Asteraceae; Asteroideae; Eupatorieae; Stevia

### References

[1] NUCLEOTIDE SEQUENCE.

TISSUE=Leaf:

PubMed=10504563 [NCBI, ExPASy, EBI, Israel, Japan]

Richman A.S., Gijzen M., Starratt A.N., Yang Z., Brandle J.E.;

"Diterpene synthesis in Stevia rebaudiana: recruitment and up-regulation of key enzymes the gibberellin biosynthetic pathway.";

Plant J. 19:411-421(1999).

## Comments

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### **Cross-references**

### Sequence databases

**EMBL** AF097311; AAD34295.1; -; mRNA.[EMBL / GenBank / DDBJ] [CoDingSequence]

### 3D structure databases

**HSSP** Q40577; 1HX9. [HSSP ENTRY / PDB]

ModBase Q9XEI0.

## 2D gel databases

SWISS-

Get region on 2D PAGE. 2DPAGE

## **Ontologies**

GO:0016829; Molecular function: lyase activity (inferred from electronic annotation GO GO:0008152; Biological process: metabolism (inferred from electronic annotation) QuickGo view.

## Family and domain databases

IPR008930; Terp cyc toroid.

IPR001906; Terp synth-like.

InterPro IPR008949; Terpenoid synth.

IPR005630; Tps metal bd.

Graphical view of domain structure.

PF01397; Terpene synth; 1.

Pfam PF03936; Terpene synth C; 1.

Pfam graphical view of domain structure.

ProDom [Domain structure / List of seq. sharing at least 1 domain]

## Other

UniRef View cluster of proteins with at least 50% / 90% / 100% identity.

## **Keywords**

None

### **Features**

None

### Sequence information

Length: **784** Molecular weight: CRC64: 284BD3972DC09051 [This is a checksum or

AA 89401 Da sequence]

380

40 10 20 30 50 60 MNLSLCIASP LLTKSNRPAA LSAIHTASTS HGGQTNPTNL IIDTTKERIQ KQFKNVEISV 70 80 90 100 110 120

SSYDTAWVAM VPSPNSPKSP CFPECLNWLI NNOLNDGSWG LVNHTHNHNH PLLKDSLSST

130 140 150 160 170 180 LACIVALKRW NVGEDQINKG LSFIESNLAS ATEKSQPSPI GFDIIFPGLL EYAKNLDINL

190 200 210 220 230 240 LSKQTDFSLM LHKRELEQKR CHSNEMDGYL AYISEGLGNL YDWNMVKKYQ MKNGSVFNSP

250 260 270 280 290 300 SATAAAFINH QNPGCLNYLN SLLDKFGNAV PTVYPHDLFI RLSMVDTIER LGISHHFRVE

310 320 330 340 350 360

IKNVLDETYR CWVERDEQIF MDVVTCALAF RLLRINGYEV SPDPLAEITN ELALKDEYAA

400

410

390

37<u>0</u>

420

LETYHASHIL	YQEDLSSGKQ	ILKSADFLKE	IISTDSNRLS	KLIHKEVENA	LKFPINTGLE	
	44 <u>0</u> YNVDNTRILK			47 <u>0</u> FYTCQSIYRE	48 <u>0</u> ELKGLERWVV	
49 <u>0</u> ENKLDQLKFA	50 <u>0</u> RQKTAYCYFS		_	53 <u>0</u> GILTTVVDDF	54 <u>0</u> FDIGGTIDEL	
55 <u>0</u> TNLIQCVEKW		57 <u>0</u> EHVRILFLAL		59 <u>0</u> AFKWQARDVT	60 <u>0</u> SHVIQTWLEL	
61 <u>0</u> MNSMLREAIW			64 <u>0</u> ALGPIVKPAI	65 <u>0</u> YFVGPKLSEE	66 <u>0</u> IVESSEYHNL	
67 <u>0</u> FKLMSTQGRL			70 <u>0</u> LHLSNGESGK	71 <u>0</u> VEEEVVEEMM	72 <u>0</u> MMIKNKRKEL	
73 <u>0</u> MKLIFEENGS			76 <u>0</u> YANDDGFTGN	77 <u>0</u> TILDTVKDII	78 <u>0</u> YNPLVLVNEN	
EEQR						Q! in F/ foi

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BLAST submission on BLAST ExPASy/SIB or at NCBI (USA)



Sequence analysis tools: ProtParam, ProtScale, Compute pl/Mw, PeptideMass, PeptideCutter, Dotlet (Java)



ScanProsite, MotifScan



Submit a homology modeling request to SWISS-MODEL

NPS@

NPSA Sequence analysis tools

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**ENZYME** 

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# NiceZyme View of ENZYME: EC 2.5.1.21

Official Name

Squalene synthase.

Alternative Name(s)

Farnesyl-diphosphate farnesyltransferase.

Farnesyltransferase.

Presqualene synthase.

Presqualene-diphosphate synthase.

Squalene synthetase.

Reaction catalysed

1. 2 farnesyl diphosphate <=> diphosphate + presqualene diphosphate

2. Presqualene diphosphate + NAD(P)H <=> squalene + diphosphate + NAD(P)(+)

Cofactor(s)

Magnesium or manganese.

Comment(s)

In the absence of NAD(P)H, presqualene diphosphate is accumulated.

Cross-references

**Biochemical** 

Pathways; map number(s)

E10

**PROSITE** PDOC00802

**BRENDA** 

2.5.1.21

PUMA2

2.5.1.21

PRIAM enzyme-

specific profiles

2.5.1.21

**Kyoto University** 

LIGAND chemical

2.5.1.21

database

**IUBMB** Enzyme Nomenclature

2.5.1.21

IntEnz

2.5.1.21

MEDLINE

Find literature relating to 2.5.1.21

MetaCyc

2.5.1.21

P53799, FDFT\_ARATH;

Q752X9, FDFT\_ASHGO; Q9HGZ6, FDFT CANGA; P37268, FDFT HUMAN;

P78589, FDFT\_CANAL; P53798, FDFT MOUSE; UniProtKB/SwissProt

Q754Z6, FDFT\_NEUCR; P53800, FDFT\_NICBE; O74165, FDFT\_PICJA;
Q02769, FDFT\_RAT; P36596, FDFT\_SCHPO; Q92459, FDFT\_USTMA;
Q9Y753, FDFT\_YARLI; P29704, FDFT\_YEAST;

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## UniProtKB/Swiss-**Prot entry** Q12051



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Note: most headings are clickable, even if they don't appear as links. They link to the user manual or other documents.

## **Entry information**

Entry name

**GGPPS YEAST** 

Primary accession number

Q12051

Secondary accession numbers

None

Integrated into Swiss-Prot on

March 21, 2006

Sequence was last modified on

November 1, 1996 (Sequence version 1)

Annotations were last modified on

June 13, 2006 (Entry version 33)

## Name and origin of the protein

Protein name

Geranylgeranyl pyrophosphate synthetase

**Synonyms** 

**GGPP** synthetase

**GGPPSase** 

Geranylgeranyl diphosphate synthase

**BET2 suppressor protein 1** Dimethylallyltranstransferase

(EC 2.5.1.1)

Includes

Geranyltranstransferase

(EC 2.5.1.10)

**Farnesyltranstransferase** 

(EC 2.5.1.29)

Gene name

Name: BTS1

OrderedLocusNames: YPL069C

From

Saccharomyces cerevisiae (Baker's yeast) [TaxID: 4932]

Taxonomy

Eukaryota; Fungi; Ascomycota; Saccharomycotina;

Saccharomycetes; Saccharomycetales; Saccharomycetaceae; Saccharomyces.

## References

[1] NUCLEOTIDE SEQUENCE [GENOMIC DNA], AND FUNCTION.

DOI=10.1074/jbc.270.37.21958; PubMed=7665600 [NCBI, ExPASy, EBI, Israel, Japan] Jiang Y., Proteau P., Poulter D., Ferro-Novick S.;

"BTS1 encodes a geranylgeranyl diphosphate synthase in Saccharomyces cerevisiae."; J. Biol. Chem. 270:21793-21799(1995).

## [2] NUCLEOTIDE SEQUENCE [LARGE SCALE GENOMIC DNA].

**STRAIN**=S288c / AB972:

PubMed=9169875 [NCBI, ExPASy, EBI, Israel, Japan]

Bussey H., Storms R.K., Ahmed A., Albermann K., Allen E., Ansorge W., Araujo R., Aparida., Barrell B.G., Badcock K., Benes V., Botstein D., Bowman S., Brueckner M., Carpenter Cherry J.M., Chung E., Churcher C.M., Coster F., 🖾 , Hani J.;

"The nucleotide sequence of Saccharomyces cerevisiae chromosome XVI."; Nature 387:103-105(1997).

131 NUCLEOTIDE SEQUENCE IGENOMIC DNA1.

## STRAIN=S288c;

Marsischky G., Rolfs A., Richardson A., Kane M., Baqui M., Taycher E., Hu Y., Vannberg Weger J., Kramer J., Moreira D., Kelley F., Zuo D., Raphael J., Hogle C., Jepson D., Williamson J., Camargo A., Gonzaga L., 🖾 , LaBaer J.;

"Creation of the YFLEX clone resource: cloning of Saccharomyces cerevisiae ORFs in the Gateway recombinational cloning system.";

Submitted (JUL-2004) to the EMBL/GenBank/DDBJ databases.

## [4] LEVEL OF PROTEIN EXPRESSION.

DOI=10.1038/nature02046; PubMed=14562106 [NCBI, ExPASy, EBI, Israel, Japan] Ghaemmaghami S., Huh W.-K., Bower K., Howson R.W., Belle A., Dephoure N., O'Shea I Weissman J.S.;

"Global analysis of protein expression in yeast.";

Nature 425:737-741(2003).

### [5] FUNCTION.

DOI=10.1111/j.1600-0854.2004.00213.x; PubMed=15296494 [NCBI, ExPASy, EBI, Israel Japan]

Shiflett S.L., Vaughn M.B., Huynh D., Kaplan J., Ward D.M.;

"Bph1p, the Saccharomyces cerevisiae homologue of CHS1/beige, functions in cell wall formation and protein sorting.";

Traffic 5:700-710(2004).

### Comments

- **FUNCTION**: Catalyzes the trans-addition of the 3 molecules of IPP onto DMAPP to forr geranylgeranyl pyrophosphate. Required for the membrane attachment of YPT1 and SI May be involved in vesicle trafficking and protein sorting.
- **CATALYTIC ACTIVITY**: Dimethylallyl diphosphate + isopentenyl diphosphate = diphos<sub>|</sub> + geranyl diphosphate.
- CATALYTIC ACTIVITY: Geranyl diphosphate + isopentenyl diphosphate = diphosphate trans,trans-farnesyl diphosphate.
- CATALYTIC ACTIVITY: Trans,trans-farnesyl diphosphate + isopentenyl diphosphate = diphosphate + geranylgeranyl diphosphate.
- PATHWAY: Isoprenoid biosynthesis.
- SUBCELLULAR LOCATION: Cytoplasm (By similarity).
- MISCELLANEOUS: Present with 2840 molecules/cell.
- SIMILARITY: Belongs to the FPP/GGPP synthetase family.

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### **Cross-references**

### Sequence databases

U39205; AAB68296.1; -; [EMBL / GenBank / DDBJ]
Genomic\_DNA. [CoDingSequence]
U31632; AAA83262.1; -; [EMBL / GenBank / DDBJ]

Genomic DNA.

[CoDingSequence]

[CoDingSequence]

**EMBL** 

AY692852; AAT92871.1; -;

[EMBL / GenBank / DDBJ]

Genomic DNA.

PIR

S60921; S60921.

3D structure databases

ModBase

Q12051.

Protein-protein interaction databases

IntAct

Q12051; -.

DIP

Q12051.

2D gel databases

SWISS-2DPAGE Get region on 2D PAGE.

Organism-specific gene databases

SGD

S000005990; BTS1.

Yeast-GFP

YPL069C.

**Ontologies** 

GO:0005739; Cellular component: mitochondrion (inferred from direct assa GO:0004311; Molecular function: farnesyltranstransferase activity (inferred

direct assav).

GO

GO:0016114; Biological process: terpenoid biosynthesis (inferred from dire

assay).

QuickGo view.

Family and domain databases

IPR000092; Polyprenyl synt.

InterPro

IPR008949; Terpenoid synth.

Graphical view of domain structure.

Pfam

PF00348; polyprenyl synt; 1.

Pfam graphical view of domain structure.

**PROSITE** 

PS00723; POLYPRENYL SYNTHET 1; 1.

ProDom

PS00444; POLYPRENYL SYNTHET 2; 1.

[Domain structure / List of seq. sharing at least 1 domain]

**BLOCKS** 

Q12051.

Genome annotation databases

Ensembl

YPL069C; Saccharomyces cerevisiae. [Contig view]

GenomeReviews U00094 GR; YPL069C.

Other

LinkHub

Q12051; -.

**ProtoNet** 

Q12051.

UniRef

View cluster of proteins with at least 50% / 90% / 100% identity.

**Keywords** 

Carotenoid biosynthesis; Complete proteome; Isoprene biosynthesis; Multifunctional enzyme; Protein transport; Transferase; Transport.

**Features** 



Feature table viewer

CHAIN	From To Len 1 335 3 169 169	-	ranyl pyropho	sphate synthet	FTId case. PRO_0000	228						
Sequence information												
Length: <b>335 AA</b> [This is the length of the unprocessed precursor]		Molecular weight: <b>38651 Da</b> [This is the MW of the unprocessed precursor]		CRC64: <b>4C7D6527FF29F157</b> is a checksum on the sequence								
1 <u>0</u> MEAKIDELIN		O 3 <u>0</u> E SLISKPYNHI	4 <u>0</u> LLKPGKNFRL	5 <u>0</u> NLIVQINRVM	6 <u>0</u> NLPKDQLAIV							
7 <u>0</u> SQIVELLHNS		<u>0</u> 9 <u>0</u> N APLRRGQTTS	10 <u>0</u> HLIFGVPSTI	11 <u>0</u> NTANYMYFRA	12 <u>0</u> MQLVSQLTTK							
13 <u>0</u> EPLYHNLITI		15 <u>0</u> R GQGLDIYWRD		17 <u>0</u> MYLNMVMNKT	18 <u>0</u> GGLFRLTLRL							
19 <u>0</u> MEALSPSSH		<u>)</u> 21 <u>0</u> L LGIIYQIRDD		23 <u>0</u> SEKGFAEDIT	24 <u>0</u> EGKLSFPIVH							
25 <u>0</u> ALNFTKTKGQ	-	<u>270</u> I LLLRTSDKDI	28 <u>0</u> KLKLIQILEF	29 <u>0</u> DTNSLAYTKN	30 <u>0</u> FINQLVNMIK	Q·						
310	32	<u>)</u> 33 <u>0</u>				in						

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NDNENKYLPD LASHSDTATN LHDELLYIID HLSEL

BLAST submission on BLAST ExPASy/SIB or at NCBI (USA)



Sequence analysis tools: ProtParam, ProtScale, Compute pl/Mw, PeptideMass, PeptideCutter, Dotlet (Java)



ScanProsite, MotifScan



Submit a homology modeling request to SWISS-MODEL



NPSA Sequence analysis tools

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